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# (54) DETERGENT COMPOSITIONS

(71) We, PROCTER & GAMBLE LIMITED, a Company organised under the laws of the United Kingdom of Hedley House, Gosforth, Newcastle-upon-Tyne, NE99 IEE, do hereby declare the invention, for which we pray that a patent may be granted to us, and the method by which it is to be performed to be particularly described in and by the following statement:—

This invention relates to compositions which protect keratinous material, such as skin or hair, from the deleterious effects of detergents or other harsh materials such as solvents, and from adverse climatic conditions. The compositions of the invention accordingly help to maintain the keratinous material in good condition.

The deleterious effects of compositions containing surfactants upon keratin are well known. These effects are caused, it is thought, by penetration of the surfactant into the keratin surface leading to "leaching out" of oils and moisturising components essential for good condition of the keratin. This penetration by surfactant and "leaching out" of essential oils also affects the ability of the keratin, particularly in the case of skin, to retain water within the tissue and this again leads to poor conditions of the keratinous material.

Many attempts have been made in the past to provide compositions for maintaining or improving the condition of skin and hair. The application of protein to skin and hair as cosmetic treatments probably antedates recorded history. Casein, in the form of milk, has been used as a time-honoured beautifier and more recently has been recommended for use in toilet soaps. U.K. Patent 1.160,485 describes the inclusion of partially degraded water soluble proteins having a gel strength of zero Bloom grams in detergent compositions and lotions for application to skin as dishwashing liquids.

German Offenlegungsschriften (Published Specifications) 2,151,739 and 2,151,740 describe certain fatty derivatives of low-molecular-weight aminolysates suitable for use in shampoos. U.K. Patent 1,122,076 describes the preparation of low-molecular-weight, alcohol-soluble protein esters suitable for use in hairspray formulations. Various low-molecular-weight polypeptides or modified polypeptides are commercially available and recommended for use in cosmetic and shampoo formulations, for instance Hydro Pro 220, and Hydro Pro 330 marketed by The Stepan Chemical Company; and Wilson X250, Wilson X1000 and Wilson Aqua Pro marketed by the Wilson Chemical Company. However, it has been found that none of these compositions are especially effective in protecting keratin from the action of harsh detergents, and this is particularly true when the proteins are incorporated in the detergent composition itself. The emolliency of these compositions can often be improved by addition of fatty or oily materials but, when used in dishwashing liquids, this usually leads to loss of foaming power or aesthetic changes which are

generally considered undesirable by consumers.

The present invention therefore provides protein-containing compositions which are particularly effective in protecting keratinous material, such as skin or hair, from the deleterious effects of detergents and other harsh materials and from adverse climatic conditions, which compositions are effective even when applied to

2	1,323,041	
	keratin in foaming detergent solutions and which result in no loss of foaming or cleaning power for detergent solutions containing them.  Accordingly, the present invention provides a detergent composition comprising	
5	(1) from 5 to 90% by weight of detergent selected from anionic, nonionic, zwitterionic and amphoteric detergents and mixtures thereof, and (2) from 0.1 to 20% by weight of a modified protein (as hereinafter defined) having an isoionic point (pI) of less than pH6 and an average molecular weight of at	5
10	least 600, in which primary amino groups of the precursor protein have been replaced by —NHYR groups, wherein Y represents carbonyl or sulphonyl and wherein R is a group containing no more than seven carbon atoms having the general formula $CH_2$ ( $CH_2$ ) <sub>q</sub> ( $CH_2$ ) <sub>q</sub> ( $OH_2$ ) in which Q and $OH_2$ are $OH_2$ or $OH_3$ or $OH_4$ is a hydrogen atom, an alkyl or alkenyl group, p is 0 or 1 and q is from 0 to (5-p).	10
15	In this specification a modified protein means a product, other than a derived protein, obtained in one or more stages by chemical modification of a precursor protein, a precursor protein being a non-enzymic protein chosen from natural, derived, synthetic or biosynthetic proteins and a derived protein being the product of hydrolytic, ammoniolytic, enzymic or thermal degradation of a protein material.	15
20	The precursor proteins suitable for use, after modification, in the compositions of the invention, may be chosen from natural, derived, synthetic or biosynthetic proteins. Natural proteins may be of either animal or vegetable origin and include simple and conjugated protein.  Typical natural proteins include intracellular proteins and globular proteins	20
25	such as those present in blood plasma and milk, as well as solubilized collagen and protein isolates from nuts, cereals etc. such as soybean isolate, peanut, protein, cotton seed protein etc. Derived proteins may be obtained from many sources, for instance by hydrolytic, ammoniolytic, thermal or enzyme degradation of globular or structural proteins such as keratin, collagen, fibrinogen, myosin, whey, egg	25
30	white, casein or vegetable proteins such as those obtained from cereals, nuts, soybean curd or the protein-rich residues from seed-oil manufacture. Preferred synthetic proteins include polylysine and unicellular proteins obtained from bacterial micro-organisms.  Protein primary amino group modification takes place primarily at lysine	30
35	groups and, desirably, the precursor protein should have at least 4 gms., preferably at least 6 gms. of lysine per 100 gms. of protein. Suitable precursor proteins in this class include the milk proteins, casein and whey, and egg white proteins (primarily ovalbumin), bacterially derived unicellular protein and soy, or derived proteins prepared therefrom. In addition, suitable precursor proteins should comprise at	35
40	least 20 gms. of aspartyl and glutamyl groups, in total, per 100 gms. of protein. Amino acid contents for a wide variety of proteins are given on page 105 of Amino Acids and Proteins by D. M. Greenberg published by Charles Thomas in 1951, which disclosure is hereby incorporated by reference.	40
45	Preferred classes of modified protein falling within the above definition are those in which R is represented by:  (1) CH <sub>2</sub> —CH(OH)—(CH <sub>2</sub> ),—H in which r is from 0 to 4, and (2) CH <sub>2</sub> —(CH <sub>2</sub> ),—H in which r is from 0 to 3.	45
50	The modified proteins of the present invention are made by modification of protein precursor side chains comprising free primary amino groups, preferably, by acylation thereof.  Methods of preparation of proteins having these functional substituents are well known in the art and the necessary preparative techniques are described in "The Chamical Medification of Proteins" by G. F. Monte and R. F. France	50
55	"The Chemical Modification of Proteins" by G. E. Means and R. E. Feaney published by Holden-Day Inc. in 1971, the disclosures of which are hereby incorporated by reference. Although the principal reactive centres are the protein side-chains comprising primary amino groups, simultaneous modification of other reactive centres such as carboxylic acid, sulphydryl, aliphatic or phenolic hydroxy, imidazole or gaunidino groups, may also occur.	55
60	In preferred embodiments of the invention, the proteins may be acylated via primary amino groups. Acylation may be performed by using the appropriate acid anhydride or N-carboxy anhydride. In the latter case, this results in acylation predominantly at amino groups. In the former case, if the acid anhydride is cyclic,	60

	the modification leads to acidic substituents which should be neutralized, for instance by esterification. Reactions analogous to acylation may also be performed. Thus, sulphonamide derivatives of proteins may be prepared, for performed the sulphonyl halides.	
5	instance, by reaction of protein primary amino groups with sulphonyl halides.  Particularly highly preferred proteins are the N-acyl and N-sulphonyl derivatives of degraded proteins, particularly of degraded casein, soyprotein and collagen (gelatin). The acyl group may be introduced by allowing an aqueous solution of the protein to stand at a temperature of 10°—20° for 1—2 hours with	5
10	n- and iso- butyric acids. The reaction may be conducted at a pH of from 6 to 9, especially from 7 to 8, using a suitable buffer to promote acylation rather than	10
15	of protein and anhydride used. Generally at least 20% of the protein primary amino groups will be acylated, preferably at least 40% and especially at least 60%; 80% or more acylation is readily achievable using a large (20 to 30 fold) excess of	15
20	anhydride. Condensation products with sulphonic acid derivatives are generally prepared via the corresponding sulphonyl chloride compounds. The sulphonyl chloride compound is conveniently obtained from the corresponding sulphonic acid by treatment with phosphorus pentachloride and is treated with an alkaline solution or dispersion of the protein at 50—100°C for, e.g. 4—5 hours, with constant addition	20
25	of aqueous alkali to maintain an alkaline pri.  Particularly preferred proteins for use in the compositions of the invention have characteristic values of molecular weight and isoionic-point pH and these will the discussed in some detail	25
30	It will be appreciate that the molecules of a protein vary widely in their size and complexity and that the molecular weight of a protein is necessarily an imprecise quantity. The molecular weight of a protein may be specified by defining the molecular weight distribution of the molecules of the protein, but it is usual to define, instead, the average molecular weight of the protein sample because it is an expectation weight of the protein sample because it is an expectation weight of the protein sample because it is an expectation weight.	. 30
35	average molecular weight which however, to the actual molecular weight average is only an approximate guide, however, to the actual molecular weight distribution of the sample. Also, it should be appreciated that the average molecular weight as measured may vary from one measuring technique to another although the differences between the results of the various techniques generally diminish towards lower molecular weights. In this specification, one method employed for determining average molecular weights of proteins (for molecular measurements of	35
40	weights greater than about 5000) makes use of viscometric measurements of buffered protein solutions. The intrinsic viscosity of a buffered protein solution is known to be primarily dependent upon the overall length of the protein coil and to be relatively independent of the nature of the sidechain and end groups of the protein. There is, therefore, a relationship between instrinsic viscosity and the average molecular weight of the protein, which may be expressed as	40
45	$[\eta] = K.M^a$ [Staudinger's Equation]	45
	where K and a are constants for a particular source of protein. It is thus straightforward to determine molecular weights from viscosity measurements, knowing K and a, and this is fully described in Macromolecular Chemistry of Gelatin, page 72, by A. Veiss, and in Biochimica et Biophysica Acta, 57, 222—9.	
50	(1961) by J. Bello, H. R. Bello, and J. R. Vinograd.  However, the viscosity method is not very accurate at molecular weights of about 5000 and below and ultracentrifuge techniques have been used in this molecular weight region. Comparison of the two techniques reveal only small	50
55	When measured by the above methods, the precursor and modified proteins of the present invention generally have molecular weights of less than about 50,000 and preferably less than about 20,000. Desirably, they fall in the range from 1,000 to	55
60	10,000 and more especially from 2,000 to 5,000.  The modified proteins are present in the compositions of the invention in an amount from 0.1% up to 20%, preferably in an amount between 1% and 10%, more preferably between 2% and 6% by weight of the compositions.  Protein molecules, having both acidic and basic side chains, are charged in both acidic and basic solutions and thus are amphoteric in nature. The pH at which	60

4	1,529,841	4
5	equal concentrations of protein anions and cations exist in solution is known as the isoionic point, and in the present invention, the isoionic point of the modified proteins must be less than 6, generally in the range 2.5 to 5.5, and preferably in the range 2.5 to 4.0. The isoionic point pH may be determined in the following manner:—	5
	Amberlite (Registered Trade Mark) acid resin (IR 120) and base resin (IR 400) are washed with several volumes of water, filtered and mixed in the ratio 0.4:1. A solution (20 mls.) of protein (3%) and urea (20% by weight) is prepared with minimum warming and allowed to cool to constant temperature. The resin mixture	
10	(8.4 g.) is added, the solution is stirred for five minutes, the mixture is filtered and the pH of the filtrate is the isoionic point pH of the protein.  A specific preparative method for acylated proteins useful in the present invention is as follows:	· 10
15	Promine F (120 g.), an edible grade soybean isolate (Promine F being a Trade Mark) was added to vigorously stirred warm water (360 mls.) to form a slurry. The mixture was heated to a slurry temperature of 90—95°C and sodium hydroxide pellets (12 g.) were added. After stirring for 4 hours, the liquid was cooled to 30°C. After cooling, acetic anhydride (60 mls.) was added slowly to the solution over a	15
20	period of about 1 hour, keeping the pH of the solution at about 7 to 8, by the addition of further sodium hydroxide, and maintaining the temperature below 20°. The solution was then stirred for a further hour, and the protein was precipitated by acidifying to pH3 and cooling. The precipitate was washed with cold acid and the protein obtained by freeze drying. It had an isoionic point pH of about 3, a molecular weight of about 3000 and contained substantially no unacetylated $\varepsilon$ -	20
25	amino groups.  A similar procedure was used to obtain higher acyl derivatives, although somewhat longer reaction times were required in these cases. The procedure was also applicable to the acylation of whole protein, such as whole casein, whey, serum albumin etc.	25
30	The optimum choice of protein for any particular composition depends to a certain extent upon the pH of the composition in use, i.e. the pH of the carrier upon application to keratin. This in-use pH may, depending upon the type of application, be the pH of the composition itself, or be the pH of an aqueous solution or dispersion of the composition at a concentration of use which may be as	30
35	In order to obtain the maximum occlusive benefit, compositions of the invention should have an in-use pH as different as possible from the pI of the modified protein. For modified proteins having a pI<6, the in-use pH is preferably greater than pH6 and desirably is greater than (pI + 2).	35
40	The in-use pH of the compositions of the invention may vary widely, of course, depending upon the purpose and manner of use of the compositions. Liquid compositions designed for shampoos are generally applied to hair in medium/high concentration aqueous solution, and the in-use pH is close to the pH of the composition itself. This may be any pH in the range, generally, from 4 to 9.	40
45	Detergent compositions such as liquid dishwashing compositions, bathing compositions and heavy-duty granular or liquid detergents are usually used in a large excess of water, and the in-use pH is the pH of an aqueous solution of the composition at a concentration generally in the range from 0.01% to 2% by weight. Builder-free detergent compositions used, for instance, as light-duty detergents	45
50	generally have an in-use pH of about 7; built heavy-duty detergents generally have an in-use pH in the alkaline range up to a pH of about 11. Soap bar compositions are applied to skin as an aqueous solution or dispersion of the soap bar ingredients at a concentration, generally in the range from 5 to 15 wt %. The pH of the soap dispersion may vary, depending upon the type of soap bar employed, from a pH of	50
55	5.5 to about 10.5.  Surfactant materials which may be used in the compositions of the invention can be selected from foaming water-soluble soap and synthetic anionic, nonionic, zwitterionic and amphoteric detergents described as below. Cationic materials may be present but preferably only in the presence of other types of detergent.	55
60	A. Anionic Soap and Non-Soap Synthetic Detergents  This class of detergents includes ordinary alkali soaps such as the sodium, potassium, ammonium, alkyl ammonium and alkylolammonium salts of higher fatty acids containing from 8 to 24 carbon atoms and preferably from 10 to 20 carbon atoms. Suitable fatty acids can be obtained from natural sources, such as plant or	60

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synthetically prepared (e.g. by the oxidation of petroleum or by hydrogenation of carbon monoxide by the Fischer Tropsch process). Resin acids are suitable, such as rosin and those resin acids in tall oil. Naphthenic acids are also suitable. Sodium and potassium soaps can be made by direct saponification of the fats and oils or by the neutralization of the free fatty acids which are prepared in a separate manufacturing process. Particularly useful are the sodium, potassium, and triethanol-ammonium salts of the mixtures of fatty acids derived from coconut oil and tallow, e.g. sodium or potassium tallow and coconut soaps.

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This class of detergents also includes water-soluble salts, particularly the alkali metal salts, of organic sulphuric reaction products having in their molecular structure an alkyl radical containing from 8 to 22 carbon atoms and a sulphonic acid or sulphuric acid ester radical. (Included in the term alkyl is the alkyl portion of higher acyl radicals). Examples of this group of synthetic detergents which form a part of the preferred compositions of the present invention are the alkali metal, e.g. sodium or potassium, alkyl sulphates, especially those obtained by sulphating the higher alcohols (8 to 18 carbon atoms) produced by reducing the glycerides of tallow or coconut oil; the alkali metal class sulphanetes of from 8 to 74 carbon tallow or coconut oil; the alkali metal clefin sulphonates of from 8 to 24 carbon atoms described, for example, in U.S. Patent 3,332,880; and the alkali metal alkyl glyceryl ether sulphonates, especially those ethers of the higher alcohols derived from tallow and coconut oil. Other anionic detergents include the alkali metal alkylbenzene sulphonates, in which the alkyl group contains from 9 to 15 carbon atoms, including those of the types described in United States Patents Nos. 2,220,099 and 2,477,383 (the alkyl radical can be a straight or branched aliphatic chain); sodium coconut oil fatty acid monoglyceride sulphates and sulphonates; salts of alkyl phenol ethylene oxide ether sulphates with 1 to 12 units of ethylene oxide are molecula and in which the alkyl radicals contain from 9 to 19 carbon oxide per molecule and in which the alkyl radicals contain from 8 to 18 carbon atoms; the reaction product of fatty acids esterified with isethionic and neutralized with sodium hydroxide where, for example, the fatty acid is oleic or derived from coconut oil; sodium or potassium salts of fatty acid amides of a methyl tauride in which the fatty acids, for example, are derived from coconut oil; sodium or potassium  $\beta$ -acetoxy- or  $\beta$ -acetamido-alkanesulphonates where the alkane has from 8 to 22 carbon atoms; and others known in the art. A number are specifically set

forth in United States Patents Nos. 2,486,921; 2,486,922; and 2,396,278 A further class of surfactants falling within this category are water-soluble salts of the organic, sulphuric acid reaction products of straight or branched, saturated, aliphatic hydrocarbon radicals having from 8 to 24, preferably from 12 to 18 carbon atoms, particularly hydrocarbon radicals of the methane series, including iso-, neo-, meso-, and n-paraffins. Water-soluble salts of C14-C16 sulphonated paraffins

are especially preferred. Other synthetic anionic detergents useful herein are alkyl ether sulphates. These materials have the formula  $R^2O(C_2H_4O)_xSO_3$  wherein  $R^2$  is alkyl or alkenyl of about 8 to 24 carbon atoms, x is 1 to 30, and M is a salt-forming cation selected from alkali metal, ammonium and dimethyl-, trimethyl-, triethyl-, dimethanol-, diethanol-, trimethanol- and triethanol- ammonium cations.

The alkyl ether sulphates are condensation products of ethylene oxide and monohydric alcohols having about 8 to 24 carbon atoms. Preferably, R2 has 14 to 18 carbon atoms. The alcohols can be derived from fats, e.g. coconut oil or tallow, or can be synthetic. Lauryl alcohol and straight-chain alcohols derived from tallow are preferred herein. Such alcohols are reacted with from 1 to 12, especially 6, molar proportions of ethylene oxide and the resulting mixture of molecular species, having, for example an average of 6 moles of ethylene oxide per mole of alcohol, is sulphated and neutralized.

The alkyl ether sulphates are described in U.S. Patent 3,332,976.

B. Nonionic Synthetic Detergents

Nonionic synthetic detergents may be broadly defined as compounds produced by the condensation of alkylene oxide groups (hydrophilic in nature) with an organic hydrophobic compound, which may be aliphatic or alkyl aromatic in

One class of nonionic synthetic detergents is commercially available under the Registered Trade Mark of 'Pluronic'. These compounds are formed by condensing ethylene oxide with a hydrophobic base having a molecular weight in the range

	1500—1800 and formed by the condensation of propylene oxide with propylene glycol.	
	Other suitable nonionic synthetic detergents include the following:	
5	1. The polyethylene oxide condensates of alkyl phenol, e.g. the condensation products of alkyl phenols having an alkyl group containing from 6 to 12 carbon atoms, in either a straight-chain or branched-chain configuration, with ethylene oxide, the ethylene oxide being present in amounts equal to 5 to 25 moles of ethylene oxide per mole of alkyl phenol. The alkyl	5
10	substituent in such compounds may be derived, for example, from polymerised propylene, diisobutylene, octene or nonene. Specific examples include nonyl phenol condensed with 20 moles of ethylene oxide, dodecyl phenol condensed with 15 moles of ethylene oxide and diiso-octyl phenol condensed with 15 moles of ethylene oxide.	10
15	2. Those derived from the condensation of ethylene oxide with a hydrophobic base formed by the product resulting from the reaction of propylene oxide and ethylene diamine. Bases having a molecular weight of the order of 2,500 to 3,000 are satisfactory and typical compounds contain from 40% to 80% polyoxyethylene by weight and have a molecular weight of from 5,000 to 11,000.	15
20	3. The condensation product of aliphatic alcohols having from 8 to 24 carbon atoms, in either straight-chain or branched-chain configuration with ethylene oxide, e.g. a coconut alcohol-ethylene oxide condensate having from 5 to 30 moles of ethylene oxide per mole of coconut alcohol, the coconut alcohol fraction having from 10 to 14 carbon atoms.	20
25	4. Other particularly preferred materials are synthetic primary alcohol ethoxylates available from the Shell Oil Company under the trade marks 'Dobanol' and 'Neodol', from Imperial Chemical Industries Ltd. under the trade mark 'Synperonic', and from Liquichimica SA under the trade name 'Lial'.	25
30	Specific examples of these synthetic alcohol ethoxylats include Dobanol 45E7, a C <sub>14</sub> —C <sub>15</sub> predominantly linear alcohol containing up to 25% 2-methyl branching condensed with an average of seven moles of ethylene oxide, Dobanol 91E8, a C <sub>5</sub> —C <sub>11</sub> predominantly linear alcohol condensed with an average of eight moles of ethylene oxide, and	30
35 .	Synperonic E-3, a condensation product of 3 moles of ethylene oxide with one mole of an alcohol mixture comprising a 2:1 ratio of C <sub>13</sub> —C <sub>13</sub> primary alcohols with 50% 2-methyl branching.  5. A detergent having the formula R <sup>3</sup> R <sup>4</sup> R <sup>3</sup> N→0 (amine oxide detergent)	35
40	wherein R <sup>3</sup> is an alkyl group containing from 10 to 28 carbon atoms, from 0 to 2 hydroxy groups and from 0 to 5 ether linkages, there being at least one moiety of R <sup>3</sup> which is an alkyl group containing from 10 to 18 carbon atoms and 0 ether linkages, and R <sup>4</sup> and R <sup>5</sup> are each selected from alkyl radicals and hydroxyalkyl radicals containing from 1 to 3 carbon atoms.  Specific examples of amine oxide detergents include:	40
45	dimethyldodecylamine oxide, dimethyltetradecylamine oxide, ethylmethyltetradecylamine oxide, cetyldimethylamine oxide, dimethylstearylamine oxide, cetylethylpropylamine oxide, diethyldodecylamine oxide, diethyltetradecylamine oxide, dipropyldodecylamine oxide, bis-(2-hydroxyethyl) dodecylamine oxide, bis-(2-hydroxyethyl)-3-dodecoxy-1-hydroxyethyl)	45
50	propylamine oxide, (2-hydroxypropyl)methylietradecylamine oxide, dimethyloleylamine oxide, dimethyl-(2-hydroxydodecyl)amine oxide, and the corresponding decyl, hexadecyl and octadecyl homologues of the above compounds.  6. A detergent having the formula	50
55	O ∥ R³—S—R⁴	
<i>33</i>	wherein R <sup>3</sup> and R <sup>4</sup> are as defined above. Specific examples of sulphoxide detergents include dodecyl methyl sulphoxide, tetradecyl methyl	55
60	sulphoxide, 3-hydroxytridecyl methyl sulphoxide, 3-methoxytridecyl methyl sulphoxide, 3-hydroxy-4-dodecoxybutyl methyl sulphoxide, octadecyl 2-hydroxyethyl sulphoxide and dodecylethyl sulphoxide.	60

5	a. From 0% to 45% of a water-soluble hydrocarbon sulphate of the general formula R <sup>2</sup> O(C <sub>2</sub> H <sub>4</sub> O) <sub>n</sub> SO <sub>3</sub> M wherein R <sup>2</sup> is a straight or branched, saturated or unsaturated aliphatic, hydrocarbon radical having from 8 to 24 carbon atoms, n is from 1 to 12; and M is an alkali or alkali earth metal, ammonium or dimethyl, trimethyl, triethyl, dimethanol, diethanol, trimethanol or triethanol ammonium cation;	5
	b. From 0% to 45% by weight of a water-soluble hydrocarbon sulphonate of the general formula R <sup>3</sup> SO <sub>3</sub> M; wherein R <sub>3</sub> is a straight or branched chain saturated or unsaturated C <sub>4</sub> —C <sub>14</sub> aliphatic hydrocarbon radical or a C <sub>4</sub> —C <sub>18</sub> straight or	40
10	branched chain alkyl benzene radical and M is as defined above; c. From 0% to 45% by weight of a water-soluble hydrocarbon sulphate of the general formula R <sup>2</sup> OSO <sub>3</sub> M; where M and R <sub>2</sub> are as defined above; d. From 0% to 10% by weight of an ammonia, monoethanol or diethanol amide of	10
15	a fatty acid having an acyl moiety of from 8 to 18 carbon atoms; e. Up to 40% by weight of the condensation product of from 3 to 25 moles of an alkylene oxide, preferably ethylene or propylene oxide, and one mole of an organic, hydrophobic compound, aliphatic or alkyl aromatic in nature, the latter having from 8 to 24 carbon atoms; and f. From 0% to 10% of a trialkyl amine oxide of formula	15
20	$R_4R_5R_4N\rightarrow O$	20
	wherein $R_4$ is an alkyl group containing from 10 to 28 carbon atoms and $R_5$ and $R_4$ are each selected from alkyl radicals and hydroxyalkyl radicals containing from 1 to 3 carbon atoms.	
25	Granular detergents conventionally contain from 5% to about 20% by weight of the composition of foaming surfactant normally a $C_{16}$ — $C_{14}$ linear alkyl benzene sulphonate, alone or in admixture with a higher alkyl sulphate such as a synthetic or naturally derived $C_{16}$ — $C_{18}$ alkyl sulphate, e.g. tallow alkyl sulphate. Optional ingredients of such mixtures are also higher alkyl ethoxysulphates such as those	25
30	disclosed in German Laid Open Applications DOS Nos. 2355940 and 2355983, both incorporated herein by reference.  Nonionic surfactants such as ethoxylated primary or secondary alcohols and/ or alkyl amides and ethanolamides may also be present at levels up to 10% of the composition.	30
35 .	Dishwashing liquid products in accordance with the invention can contain up to 45% surfactant by weight of the composition. Typical formulations include 15%—30% of a linear alkylbenzene sulphonate and 15%—30% of a C <sub>10</sub> —C <sub>18</sub> linear alkyl ethoxy sulphate containing an average of 1—6 ethoxy groups. Another preferred formulation includes 5%—15% of a C <sub>14</sub> —C <sub>16</sub> paraffin sulphonate, 5—15%	35
40	of $C_{12}$ alkyl triethoxysulphate and 1%—10% of each of $C_{12}$ amine oxide, coconut alcohol hexaethoxylate and lauric diethanolamide. Another dishwashing liquid having mild properties to skin includes from about 15% to about 25% of a $C_{12}$ — $C_{14}$ alkyl ether sulphate containing an average of from about 3 to about 12 ethylene oxide groups, up to about 10% of a $C_{12}$ — $C_{14}$ alkyl sulphate, from about 1% to about 5% of a $C_{12}$ — $C_{14}$ alkyl glyceryl ether sulphonate and from about 2% to about 6% of	40
45	a $C_{12}$ — $C_{14}$ alkyl dilower alkyl amine oxide.  The invention has been found to be of particular value in liquid detergent compositions comprising a $C_8$ — $C_{18}$ linear or branched alkyl benzene sulphonate and an auxiliary natural or synthetic anionic, nonionic, amphoteric or zwitterionic surfactant in which the weight ratio of the alkyl benzene sulphonate to the auxiliary	45
50	surfactant is in the range from 1.5:1 to 10:1, preferably from 1.8:1 to 5:1 and most preferably from 2.5:1 to 4:1. In such compositions the ratio of alkyl benzene sulphonate plus auxiliary surfactant to the modified protein should be in the range of 4:1 to 100:1. Preferred examples of the auxiliary surfactants are the water-soluble	50
55	alkyl ether sulphates, particularly the $C_{12}$ — $C_{16}$ alkyl ether sulphates containing from 1 to 12 moles of ethylene oxide per mole of alkyl ether sulphate.  A further preferred liquid detergent composition contains from 10%—25% by weight of a water-soluble saturated hydrocarbon sulphonate, from 1%—10% by weight of an alkyl ether sulphate and from 1%—15% by weight of a water-soluble solvent such as a lower alkanol, in addition to the usual minor ingredients such as	55
60	perfume, colour, buffers, anti-tarnish agents etc.  The liquid detergent or gel compositions of the invention generally comprise a carrier based upon water and/or a water-soluble solvent. Suitable solvents include	60

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	C <sub>2-6</sub> mono and di-alcohols, e.g. ethanol, butanol, methyl propanol-1 and -2, amylol or pentanol, butanediol, toluol, benzyl carbinol, ethyleneglycol monobutyl ether, propyleneglycol propyl ether and diethyleneglycol dimethyl ether. They are	
5	generally present in amounts up to 13% by weight of the second of the se	5
	fluorescers, tarnish inhibitors, bactericides, hydrophobic oily materials and	
10	alkylaryl sulphonates such as south all position at levels of up to 10% sulphonate, benzene sulphonate and cumene sulphonate at levels of up to 10% sulphonate, benzene sulphonate and cumene sulphonate at levels of up to 10% sulphonate.	10
	hydrotropes such as methanot, ethanot, propulse in the composition. levels of 1—15%, normally 5—15% by weight of the composition.	15
15	Hydrophobic oily materials suitable for use in the hydrophobic oily materials suitable for use in the hydrophobic oily materials such as stearyl, myristyl and cetyl alcohols; fatty and carnauba wax; fatty alcohols such as stearyl, myristyl and cetyl alcohols; fatty and carnauba wax; fatty alcohols such as stearyl myristate and glyceryl monostearate; esters and partial esters such as isopropyl myristate and glyceryl monostearate; fatty acids such as stearic acid; lanolin and cholesterol derivatives; and silicone oils. The fatty acids such as stearic acid; lanolin and cholesterol derivatives; and silicone oils. The	
<b>2</b> 0 .	fatty acids such as stearic acid; ranolin and choicested derivative designed to enhance compositions of the invention may also comprise components designed to enhance the moisturizing effectiveness of the compositions. Suitable components include the moisturizing effectiveness of the compositions and 2 to 3 hydroxy groups,	20
25	for example 1,4 butanedio, 1,2-propylene gryest and gryendine, pyrrolidone or components include urea or urea derivatives such as guantidine, pyrrolidone or allantoin.	25
	depressants, bleaches, anti-redeposition agents, chaymos, observed of granular activators, fluorescers, builders and other normal components of granular determent compositions. Solid compositions in bar form may also contain additives	. 30
30	such as fatty acids, saits, skin creams and ons.	
35	Conditioning performance was measured in both in-vitro and in-vivo tests, a high degree of correlation between the two test methods being found. The in-vitro test (called the calf-skin occlusivity test) was based upon the rate of water transpiration through a sample of calf-skin brought into contact with a 0.15% aqueous solution of a detergent composition (at 18° hardness) containing the protein. The occlusivity of the protein was measured as the percentage reduction in the rate of water transpiration for the proteinaceous surfactant solution compared	35
40	of normal housewives in a multi-product test; hands were balanced for right hand/ left hand differences, so that there were 32 hands per product, 16 right and 16 left.	40
45	week. Treatment solutions were replenished every 10 minutes. Hands were withdrawn and reimmersed in the solution every 2 minutes. Hands were graded withdrawn and reimmersed in the solution every 2 minutes. Hands were graded withdrawn and reimmersed in the solution every 2 minutes. Hands were graded withdrawn and reimmersed in the solution every 2 minutes.	45
50	here, on a scale in which an 0.15% aqueous solution of a standard detergent was assigned HIT grades of 0, and a 1 mg/cm <sup>2</sup> application of hand-care lotion was	50
5.5	A second <i>in-vivo</i> test was also used which compares the rate of water loss through skin, specifically human forearm skin, under controlled conditions before and after treatment with a test solution. The lower the rate of water loss, the greater the ability of the skin to bind moisture and hence the greater the resistance to	55
55	chapping, cracking etc.  A test area of skin was treated with a compound (Poldine methyl sulphate) that  A test area of skin was treated area was then swent with dry nitrogen and the	
60	inhibits sweating and the treated alca was this water content of the gas assayed before and after passage over the skin. The Test water content of the gas assayed before and after passage over the skin. The Test area was immersed in an 0.15% product solution for 10 minutes at 45°C (Water hardness = 18°H), dried, and allowed to equilibrate for one hour before a further reading was obtained of the water content of a dry nitrogen sream passed over the treated skin surface. The change in water loss before and after treatment expressed	
	Could brill be a second of the	•

10		-,0-25,011				
	conditioning power of	the water loss before to the particular protein, abodiments of the inven				
5	EXAMPLE I.  Two liquid detergent compositions (identified as Example I and Standard I) were made up and tested using the <i>in-vivo</i> test method set out previously for assessing the water loss through skin. The compositions had the following ingredients:					
10	Composition		Example I	Standard I	10	
	Ammonium C <sub>12</sub> —C <sub>14</sub> li sulphonate	near alkyl benzene	18.4	18.4		
	Sodium C <sub>12</sub> —C <sub>14</sub> linear sulphate	triethoxy alkyl	18.4	18.4		
15	C12 monoethanolamide	:	2.0	4.5	15	
	Industrial Methylated	Spirits	13.0	11.0		
20	Single Cell Protein* ac hydrolysate in which ac groups was substantiall	cetylation of the amino				
20	Estimated pI = 3—4		3.6	_	20	
	Water		to 100	to 100		
	*Toprina G, a protei supplied by British Pe	in derived from yeast of troleum Ltd.	cultured on purif	ied alkines and		
25	In the above-me follows:—	ntioned in-vivo test the	two composition	ns performed as	25	
•	Standard I	+ 20 ± 10.5%	i.e. an increase loss as a res treatment			
	Example I	$-27 \pm 11\%$				
30	It can be seen that significantly better that	at Example I (acetylated, in the standard composit	hydrolysed single ion which is prote	e cell protein) is ein-free.	30	
35	EXAMPLES II and III.  Three liquid detergent compositions identified as Examples II and III and Standard II were made up and tested using the <i>in-vivo</i> test method employed in Example I. Each composition had the Base formulation of Example I and contained 3% by weight protein as follows:—					
	Standard II	Alkali-hydrolysed W	hey protein (pI = 5	5.0)		
40	Example II	Alkali-hydrolysed ac pI = 3—4, in whic groups were acetylat	h substantially a	otein; estimated ll of the amino	40	
•	Example III	Alkali-hydrolysed ac substantially all the a	etylated soybean mino groups were	protein in which acetylated.		
	In the in-vivo test,	the compositions had th	e following perfo	rmance:—		
	Standard II	+ 3 ± 8%				
45	Example II	$-7.5 \pm 8\%$			45	
	Example III	$-15.5 \pm 5\%$				

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it can be seen that Examples II and III containing proteins modified in accordance with the invention provide an occlusive benefit.

EXAMPLE IV.

The composition of Example IV was the same as Example I except that the protein was prepared by hydrolysis of soyprotein isolate followed by acylation with acetic anhydride, as described above. The N-acetylated protein had an isoionic point pH of 3, a molecular weight of about 3000 and a degree of N-acylation of about 100% of  $\epsilon$ -amino acids. The hand immersion test value for Example IV was 48 compared with a value for Standard I of 0.

EXAMPLE V.

A dishwashing liquid which is mild to skin has the following compositions:— 10

	•	Parts by wt.	
	Coconut alcohol-ethylene oxide (12) sulfate ammonium salt	18.75	
15	Coconut alcohol sulfate, ammonium salt	5.8	15
	Sodium alkyl glyceryl ether sulfonate (where the alkyl is derived from "Middle-cut" coconut alcohols and has the following approximate compositions:	·	
	2% C <sub>10</sub> ; 66% C <sub>12</sub> ; 23% C <sub>14</sub> ; 9% C <sub>16</sub>	4.0	
20	Coconut alkyl dimethyl amine oxide (wherein the coconut is middle cut)	5.0	. 20
	Potassium toluene sulfate	0.5	
	Potassium chloride	2.5	
	Citric Acid	0.1	
25	Hydrogen chloride	0.81	25
	Ammonium Xylene sulfate	5.0	
	Ethanol	8.75	
30	Protein — N-acetylated alkali degraded Promine F; isoionic point 3.0; molecular weight 3000	4.0	30
	Water	to 100	•

Similar results are obtained when the protein is replaced by N-acetylated whole casein.

## EXAMPLES VI-IX.

Granular detergent compositions were prepared with the following compositions:—

5		Example VI	Example VII	Example VIII	Example IX	5
	Sodium linear dodecyl sulphonate		6	6		
	Sodium tallow alkyl sulphate	<u> </u>	4	4	_	
10	Sodium soap (90 Tallow/ 10 Coconut)	0.5	2.5	2.5	0.5	10
	Coconut monoethanolamide	_	1.5	1.5	. –	
	Tallow monoethanolamide	0.25		_	0.25	
15	3-(N,N-dimethyl-N-C <sub>14.8</sub> -alkylammonio)-2-hydroxy-propane-1-sulphonate)	6		-3:	6	15
	Tergitol (Registered Trade Mark) 15—S—9	6	_	,	6	
20	Sodium tripolyphosphate	2	33	10	2	20
	Sodium silicate	37.5	7	7	7 .	
	Sodium carboxymethyl cellulose	0.5	0.5	0.5	0.5	
	Sodium sulphate	34	8	31	30	
25	Sodium perborate .	25	25	8	25	25
	Sodium chloride	1.5		_	1.5	
	Protease	Present	Present	Present	Present	
	Sodium EDTA	0.25	0.25	0.25	0.25	
	Perfume/brighteners	0.6	0.6	0.6	0.6	
30	Inert impurities	3.9	0.65	0.65	3.9	30
	Moisture	7	7	7	7	
	Protein	4	4	4	4	

The protein was an N-acetylated alkali degraded soyprotein having an isoionic point of 3 and a molecular weight of about 3000.

35

EXAMPLES X AND XI.

The compositions and performance of two liquid detergent products falling within the scope of the present invention, are given below. The conditioning effectiveness of these compositions, measured as the percentage reduction in the rate of water transpiration in both *in-vitro* and *in-vivo* tests, is also recorded.

40

35

13	1,529,841			13
	Composition	Exa	mples	
	Competition	x	XI	
	Ammonium linear C <sub>12</sub> —C <sub>14</sub> alkyl benzene sulphonate	27.6	27.6	
5	Sodium linear C <sub>12</sub> —C <sub>14</sub> alcohol sulphate including 3 ethylene oxide moieties	9.2	9.2	5
	Lauric monoethanolamide	2.0	2.0	
	Industrial Methylated Spirits	11.0	11.0	
	Magnesium chloride	2.1	2.1	
10	N-acetylated base-hydrolysed soyprotein  — Mol. wt. 5,000  — pI 3.0	3.0	2.0	10
	Water	to 100	to 100	
	Performance			
15	In-vitro (Percentage reduction of water transpiration) In-vivo ( — do — )	16	12 17	15
10	EXAMPLE XII.  The following liquid detergent composition was n	nade up:—		
			Wt%	•
	Sodium C <sub>14</sub> alkane sulphonate		14.4	
20	Sodium coconut alkyl triethoxy sulphate		3.6	20
	Ethyl Alcohol	•	5.0	
	Colour Perfume etc.	€ .	1.0	
	Water		To 100	
25	Acetylated hydrolysed soyprotein of pI 3.0 was the formulation at a level of 4% of the composition a tested <i>in-vivo</i> for the rate of water loss through for described herein.			25
	Results were as follows:—			
	Base formulation $+ 3 \pm 7\%$			
<b>3</b> 0	Base formulation with acetylated soyprotein $-9 \pm 7\%$			30
	The composition in accordance with the inventi a reduction in water loss from the skin whilst the give the same benefit.	on can be base compo	seen to provide osition does not	
35	WHAT WE CLAIM IS:— 1. A detergent composition comprising			35
40	<ol> <li>from 5 to 90% by weight of detergent select zwitterionic and amphoteric detergents and r</li> <li>from 0.1 to 20% by weight of a modified prote having an isoionic point (pI) of less than pH weight of at least 600, in which primary am</li> </ol>	in (as herei	nbefore defined) erage molecular	40

where R<sup>2</sup> and M are as defined above,

(d) from 0% to 10% by weight of an ammonia, monoethanol or diethanol amide of a fatty acid having an acyl moiety of from 8 to 18 carbon atoms,

(e) from 0% to 40% by weight of the condensation product of from 3 to 25 moles of an alkylene oxide per mole of an organic, hydrophobic, aliphatic or alkyl aromatic compound having 8 to 24 carbon atoms, and

(f) from 0% to 10% of a trialkyl amine oxide of formula

55

### $R_4R_5R_6N\rightarrow 0$

wherein  $R_4$  is an alkyl group containing from 10—28 carbon atoms and  $R_5$  and  $R_4$  are each selected from alkyl radicals and hydroxyalkyl radicals containing from 1—3 carbon atoms.

50

55

	13. A composition according to any preceding claim in liquid form comprising water and/or a water-soluble solvent.	
5	branched $C_s$ — $C_{18}$ alkyl benzene sulphonate and an auxiliary natural or synthetic anionic, nonionic or zwitterionic surfactant in which the weight ratio of the alkyl benzene sulphonate to the auxiliary surfactant is in the range from 1.5:1 to 10:1 and in which the weight ratio of total surfactant to modified protein is in the range from	5
10	4:1 to 100:1.  15. A composition according to Claim 14 wherein the weight ratio of the alkyl benzene sulphonate to the auxiliary surfactant is in the range from 1.8:1 to 5:1.  16. An aqueous liquid detergent composition according to either Claim 12 or Claim 13, comprising from 10%—25% of alkane sulphonate, from 1—10% of an	10
15.	alkyl ethoxy sulphate and from 1-15% of a water-soluble solvent.  17. A composition according to either Claim 12 or Claim 13 consisting essentially of from about 15-25% C <sub>12</sub> -C <sub>14</sub> alkyl ether sulphate containing an average of from three to twelve ethylene oxide groups, from 0% to 10% by weight of a C <sub>12</sub> -C <sub>14</sub> alkyl sulphate, from 1% to 5% of a C <sub>12</sub> -C <sub>14</sub> alkyl glyceryl ether	15
20	sulphonate and from 2% to 6% of a C <sub>12</sub> —C <sub>14</sub> alkyl dilower alkyl amine oxide.  18. A composition according to any of Claims 1 to 11 in bar form.  19. A composition according to Claim 1 substantially as described with reference to any one of Examples I to XII.	20

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